



#### Research Article

# Molecular and morphological evidence supports the resurrection of *Chrysosplenium guangxiense* H.G.Ye & Gui C.Zhang (Saxifragaceae)

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#### Abstract

Chrysosplenium guangxiense H.G.Ye & Gui C.Zhang was first described as a new species in 1994 but later synonymized in the Flora of China treatment with C. glossophyllum H.Hara. Plastid genomes and nrDNA sequences were used to infer the phylogenetic relationships of selected taxa in *Chrysosplenium*. Our phylogenetic analyses revealed that C. guangxiense belongs to sect. Alternifolia, is closely related to Chrysosplenium Hydrocotylifolium H.Lév. & Vaniot but distant from C. glossophyllum. Morphologically, C. guangxiense could be easily distinguished from C. glossophyllum by having robust rhizomes, basal leaves with a long cuneate base and fewer teeth in the margin, curled sepal margins, and red, larger seeds. It could also be easily distinguished from C. hydrocotylifolium by possessing long elliptic leaves and a long cuneate leaf base. Along with the phylogenetic studies, the complete plastid genome of C. guangxiense was also reported. The plastid genome was 154,004 bp in length and comprised two inverted repeats (IRs) of 28,120 bp, separated by a large single-copy of 80,646 bp and a small single-copy of 17,118 bp. A total of 111 functional genes were discovered, comprising 78 protein-coding genes, 29 tRNA genes, and four rRNA genes. Based on assessment of morphological and molecular data Chrysosplenium guangxiense H.G.Ye & Gui C.Zhang is resurrected from C. glossophyllum H.Hara at species level. A global conservation assessment classifies C. guangxiense as Vulnerable (VU).

Key words: Chrysosplenium, phylogeny, plastid genome, Saxifragaceae, taxonomy

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#### Introduction

Chrysosplenium L. (Saxifragaceae) comprises more than 70 species of perennial herbs (Kim et al. 2019; Fu et al. 2020, 2021). Chrysosplenium is distributed throughout Asia, America and Europe (Pan and Ohba 2001; Soltis 2007). The latest checklist of Chinese Chrysosplenium included 35 species (Pan and Ohba 2001). Their earlier revisions classified the genus into two subgenera (subg. Chrysosplenium and subg. Gamosplenium) based on leaf arrangement (Pan 1986a, b). This

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character was also considered by Hara, who divided *Chrysosplenium* into two sections, namely sect. *Alternifolia* and sect. *Oppositifolia* (Hara 1957). The following molecular analyses (Soltis et al. 2001; Fu et al. 2021) demonstrated that these two subgenera/sections are monophyletic and sister to each other, further confirming that leaf arrangement is a good indicator of the relationships within the genus. However, a recent systematic study based on a complete chloroplast genome and nrDNA data challenged this relationship as their results recovered an additional clade composed of two species with alternate leaves (as members of sect. *Alternifolia*). The newly defined clade was recognized as a basal clade sister to the rest of the species of *Chrysosplenium* (Yang et al. 2023).

Chrysosplenium guangxiense H.G.Ye & Gui C.Zhang was first described as a new species in 1994 by having ovate-elliptic leaves, an acuminate apex, a cuneate base, fewer dentate margins, and a depressed sepal apex, enabling it to be distinguished from its similar species, *C. glossophyllum* H.Hara (Ye and Zhang 1994). Subsequently, it appeared as a synonym of the latter in *Flora of China* without additional explanation (Pan and Ohba 2001). We assumed that the authors considered these differences were insufficient to distinguish them. It is possible that the scarcity of *C. glossophyllum* species in China, with its only population in Sichuan Province, may have led to this misinterpretation. Molecular data, however, could provide a means to confirm the systematic position of morphological similarities and to evaluate the phylogenetic informativeness of morphological characters (Scotland et al. 2003).

In 2019, we conducted an extensive investigation in Tianlin County, Baise City, Guangxi, China, the type locality of *Chrysosplenium guangxiense*. We collected a plant of *Chrysosplenium*, which was then confirmed as *C. guangxiense*. Following a thorough literature survey (Hara 1957; Pan 1992; Pan and Ohba 2001; Liu et al. 2016; Wei 2018; Kim et al. 2019; Fu et al. 2020, 2021; Wei et al. 2022), along with the molecular evidence, it was confirmed that *C. guangxiense* is a different species from *C. glossophyllum*.

#### **Materials and methods**

#### Morphology observations and conservation assessments

All morphological characters were studied based on the material from field and herbarium specimens using a dissecting microscope (SMZ171, Motic, China). For seed morphology, we also undertook scanning electron microscope (SEM) observations; seeds were collected from the field and dried with silica gel. The pre-treatments, including impurity removal, air-drying, and gold-coating, were performed following Fu et al. (2020). Observations and photographs were taken under a Hitachi SU8010 scanning electron micrograph. At least 15 seeds were used to determine their size and ornamentation. A conservation assessment was undertaken following the IUCN (2019).

### Genomic DNA extraction, sequencing, plastid genome and nrDNA assembly and annotation

The genomic DNA was extracted using the modified CTAB method (Doyle and Doyle 1987). The short-insertion library (300 bp) was constructed and then sequenced to obtain 2×150 bp paired-end data using the Illumina NovaSeq

platform at Majorbio Company (Shanghai, China). The raw data was filtered through Trimmomatic v. 0.39 (Bolger et al. 2014) to obtain clean data, and then the clean data were quality-controlled using FastQC v. 0.11.9 (Simon 2020). The complete plastid genome and nrDNA sequence were assembled using GetOrganelle v. 1.7.5 (Jin et al. 2020), and annotation was performed using CPGA-VAS2 (Shi et al. 2019) and PGA (Qu et al. 2019).

#### Phylogenetic analysis

To confirm the phylogenetic placement of *Chrysosplenium guangxiense*, we undertook phylogenetic studies using the chloroplast (CP) genomes and nrDNA sequences obtained in a previous study (Yang et al. 2023). Forty-seven species of *Chrysosplenium* as in-group, and two species from other genera in Saxifragaceae and *Itea chinensis* Hook. & Arn. from Iteaceae as an out-group were sampled. The species names and GenBank accession numbers are listed in Table 1.

The chloroplast protein-coding genes (cpPCGs) were extracted from the CP genome using PhyloSuite v.1.2.3 (Zhang et al. 2020). These cpPCGs and nrDNA sequences were aligned by MAFFT v. 7.4 (Katoh and Standley 2013), and concatenated using PhyloSuite v.1.2.3 (Zhang et al. 2020) to form the cpPCGs+nrDNA matrix. The phylogenetic analyses of *Chrysosplenium* based on cpPCGs, nrDNA and cpPCGs+nrDNA matrices were performed using maximum likelihood (ML) and Bayesian inference (BI), respectively. The ML analyses were conducted using IQ-TREE v. 2.1.2 (Nguyen et al. 2015) with 1,000 bootstrap replicates and the default ModelFinder (Kalyaanamoorthy et al. 2017) to find GTR+F+I+G4 as the best-fit substitution model. Tree visualization was achieved in Figtree v. 1.4.3 (http://tree.bio.ed.ac.uk/software/figtree/). For BI analysis, MrBayes v. 3.2.6 (Ronquist et al. 2012) was employed to obtain a maximum clade credibility (MCC) tree. BI analysis was performed using one million generations, two runs, 25% trees discarded as burn-in, and trees sampled every 1,000 generations (1,000 trees sampled in total) with the GTR model.

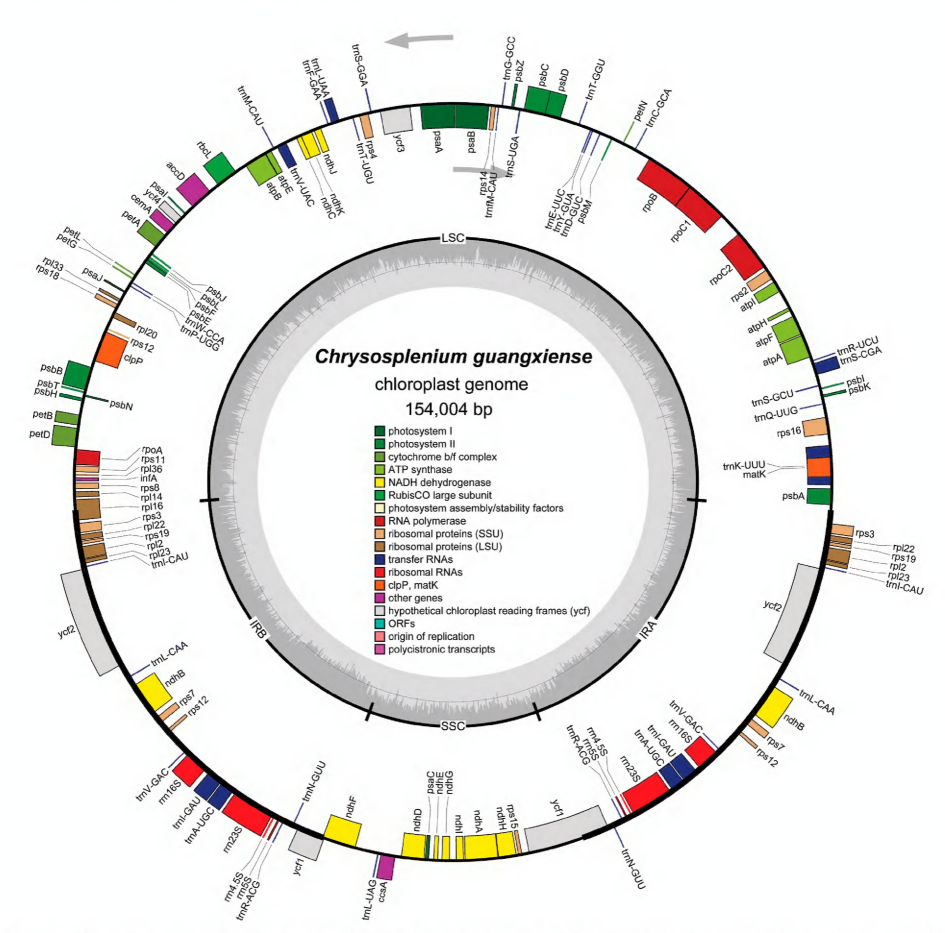
#### Results

#### Characteristics of the complete chloroplast genome

The CP genome of *Chrysosplenium guangxiense* comprised 154,004 bp (Fig. 1). The characteristics and statistics of the CP genome are summarized in Tables 4, 5.

#### Molecular phylogenetic studies

The cpPCGs matrix length was 71,919 bp, including 6,392 parsimony informative sites, 13,645 variable sites, and 55,865 conserved sites. The nrDNA matrix was 6,738 bp in length, with 765 parsimony informative sites, 1,200 variable sites, and 5,231 conserved sites. The cpPCGs+nrDNA matrix was 78,657 bp in length, with 7,157 parsimony informative sites, 14,845 variable sites, and 61,096 conserved sites. The phylogenetic tree of the cpPCGs matrix exhibited high confidence, while the phylogenetic tree of the nrDNA matrix had some branches with low support, and was significantly different from the former (Suppl. materials 1, 2). However, *Chrysosplenium guangxiense* was consistently related to



**Figure 1**. Plastid genome map of *Chrysosplenium guangxiense*. The thick lines on the outer complete circle identify the inverted repeat regions (IRa and IRb). The innermost track of the plastome shows the GC content. Genes on the outside and inside of the map are transcribed in clockwise and counter directions, respectively.

C. hydrocotylifolium H.Lév. & Vaniot in both chloroplast and nuclear gene trees (Suppl. materials 1, 2). The phylogenetic tree of the cpPCGs+nrDNA matrix received a higher confidence value compared to trees generated from subsets (cpPCGs and nrDNA). Topologies obtained from BI and ML methods were congruent and showed that Chrysosplenium species clustered in a strongly supported clade (BS = 100%, PP = 1) which was further divided into three well-supported clades (defined as A-C clades; Fig. 2). Chrysosplenium guangxiense was recognized as a member of clade B and fell in its basal clade (BS = 100%, PP = 1; Fig. 2), which also included C. macrophyllum Oliv., C. zhangjiajieense X.L.Yu, Hui Zhou & D.S.Zhou, C. hydrocotylifolium, C. flagelliferum F.Schmidt, and C. zhouzhiense Hong Liu. Of these, C. guangxiense was most closely related to C. hydrocotylifolium (BS = 100%, PP = 1; Fig. 2). Although C. glossophyllum was also a member of clade B, it fell into a much more distant clade from C. guangxiense (Fig. 2).

**Table 1.** Species names and GenBank accession numbers of plastid genomes and nrDNA sequence used in this study (\* newly generated sequences).

Species	Location	Voucher specimens	Herbarium	Plastid GenBank number	nrDNA GenBank number
Chrysosplenium album Maxim.	Nikkou-shi, Japan	HSN09815	HSN	OK336556	OP154009
Chrysosplenium aureobracteatum Y.I.Kim & Y.D.Kim	Gangwon Province, South Korea	KYI-2009032	(Kim et al. 2018)	MG878089	MK989509
Chrysosplenium biondianum Engl.	Shanxi, China	HZ2017050107362	HSN	OK336542	OP154015
Chrysosplenium carnosum Hook.f. et Thoms.	Sichuan, China	HSN013113	HSN	OK336564	OP154016
Chrysosplenium davidianum Decne. ex Maxim.	Sichuan, China	HSN06442	HSN	OK336537	OP154017
Chrysosplenium delavayi Franch.	Sangzhi, Hunan, China	SZ2016080907105	HSN	OK336539	OP154018
Chrysosplenium dubium J. Gayex DC.	Georgia	P03_WF11	(Folk et al. 2019)	-	OP154019
Chrysosplenium echinus Maxim.	Nikkou-shi, Japan	HSN09817	HSN	OK336557	OP154020
Chrysosplenium fauriae Franch.	Nikkou-shi, Japan	HSN09823	HSN	OK336561	OP154021
Chrysosplenium flagelliferum Fr. Schmidt.	Nikkou-shi, Japan	HSN09816	HSN	OK336541	OP154022
Chrysosplenium forrestii Diels	Nikkou-shi, Japan	HSN7797	HSN	OK336565	OP154024
Chrysosplenium giraldianum Engl.	Sichuan, China	JZ2018042507981	HSN	OK336548	OP154025
Chrysosplenium glossophyllum H. Hara	Sichuan, China	QCS2017102608035	HSN	OK336544	OP154026
Chrysosplenium grayanum Maxim.	Nikkou-shi, Japan	HSN09810	HSN	OK336555	OP154027
Chrysosplenium griffithii Hook.f. et Thoms.	Shanxi, China	HSN7760	HSN	OK336547	OP154028
Chrysosplenium guangxiense H.G.Ye & Gui C.Zhang	Guangxi, China	HSN13356	HSN	OP093635*	OR941245*
Chrysosplenium henryi Franch.	Sangzhi, Hunan, China	HSN7505	HSN	OK336532	OP154030
Chrysosplenium hydrocotylifolium H. Lév. & Vaniot	Hubei, China	HSN09188	HSN	OK336540	OP154031
Chrysosplenium japonicum (Maxim.) Makino	Zhejiang, China	HSN7909	HSN	OK336554	OP154032
Chrysosplenium kamtschaticum Fisch. ex Seringe	Shimane-ken, Japan	DG2019032310004	HSN	MT371065	OP154033
Chrysosplenium kiotense Ohwi.	Nikkou-shi, Japan	HSN09818	HSN	OK336558	OP154034
Chrysosplenium lanuginosum Hook.f. et Thoms.	Anhui, China	BD2017030507343	HSN	OK336534	OP154035
Chrysosplenium lectus-cochleae Kitagawa	Jilin, China	HSN7379	HSN	OK336550	OP154036
Chrysosplenium macrophyllum Oliv.	Hubei, China	BD2017030507344	HSN	MK973001	OP154037
Chrysosplenium macrospermum Y.I.Kim & Y.D.Kim	Jilin, China	CBS2016062406656	HSN	OK336562	OP154038
Chrysosplenium macrostemon Maxim. ex Franch. et Sav.	Nikkou-shi, Japan	HSN09820	HSN	OK336560	OP154039
Chrysosplenium microspermum Franch.	Jinfo Mountain, Chongqing, China	HSNTG025	HSN	OK336546	OP154040
Chrysosplenium nepalense D.Don	Tengchong, Yunnan, China	GLGH20170607375	HSN	OK336535	OP154043
Chrysosplenium nudicaule Bunge	Gansu, China	HSN07772	HSN	MZ424445	OP154044
Chrysosplenium oppositifolium L.	Wales, UK	BGN_RN_W	(Folk et al. 2019)	OR397749	OP154057
Chrysosplenium pilosum Maxim.	Nikkou-shi, Japan	HSN09819	HSN	OK336559	OP154045
Chrysosplenium valdepilosum (Ohwi) S.H.Kang & J.W.Han	Jilin, China	HSN09819	HSN	OR397753	OP154046
Chrysosplenium qinlingense Z.P.Jien ex J.T.Pan	Sichuan, China	HSN7980	HSN	OK336549	OP154047
Chrysosplenium ramosum Maxim.	Jilin, China	SJH2017052107372	HSN	MK973002	OP154048
Chrysosplenium sedakowii Turcz.	Irkutsk, Russia	P02_WC8	(Folk et al. 2019)	_	OP154049

Species	Location	Voucher specimens	Herbarium	Plastid GenBank number	nrDNA GenBank number
Chrysosplenium serreanum HandMazz.	Jilin, China	SJH2017052107371	HSN	OK336538	OP154050
Chrysosplenium sinicum Maxim.	Hunan, China	TPS2017042407504	HSN	MT362051	OP154051
Chrysosplenium taibaishanense J.T.Pan	Shanxi, China	HSN7761	HSN	OK336552	OP154052
Chrysosplenium uniflorum Maxim.	Tibet, China	HSN7380	HSN	OK336533	OP154053
Chrysosplenium zhouzhiense Hong Liu	Shanxi, China	HSN13356	HSN	OK336551	OP154055
Chrysosplenium alternifolium L.	Shimane-ken, Japan	DG2019032310003	HSN	OK336545	OP154010
Chrysosplenium tetrandrum (N. Lund) Th. Fries	Nunavut, Canada	Brysting_01-065_CAN	CAN	OR397750	OP154052
Chrysosplenium wrightii Franch. & Sav.	Yukon, Canada	Bennett_08-125_CAN	CAN	OR397751	OP154059
Chrysosplenium valdivicum Hook.	Chile	P04_WG8	HSN	OR397752	OP154060
Chrysosplenium zhangjiajieense X.L.Yu, Hui Zhou & D.S.Zhou	Hunan, China	ZJ2016031506369	HSN	OK336563	OP154054
Peltoboykinia tellimoides (Maxim.) Hara	Henan, China	PT210814	(Yang et al. 2022)	MZ779205	JQ895246
Saxifraga stolonifera Curt.	Hubei, China	S313	(Chen et al. 2022)	NC_037882	MK092506
Itea chinensis C.K.Schneider	Hunan, China	S371	-	NC_037884	MG730867

#### Morphological observations

A suite of morphological characters including rhizome size, leaf shape, leaf margin dentate, sepal apex shape, and seed size of *Chrysosplenium guangxiense* and *C. glossophyllum* was consulted or observed. *Chrysosplenium guangxiense* had a robust rhizome, basal leaves with a long cuneate base and fewer teeth in the margin, curled sepal margins, and red, larger seeds that make it easily distinguished from *C. glossophyllum* (Table 2). Considering the phylogenetic results, a morphological comparison between *C. guangxiense* and *C. hydrocotylifolium* was also conducted. *Chrysosplenium guangxiense* had long elliptic leaves and a long cuneate leaf base, which can be easily distinguished from *C. hydrocotylifolium* (Table 3).

Table 2. Diagnostic comparison of Chrysosplenium guangxiense and C. glossophyllum.

Characters	C. guangxiense	C. glossophyllum	
Rhizome	Rhizome thick, crossed and nodular	absent	
Basal leaves	base long cuneate, margin 10–20-crenate	base rounded to subcordate; margin 20–36-crenate	
Sepals	margin curl	margin uncurl	
Seed	red, 0.59-0.85 × 0.48-0.63 mm black, 0.50 × 0.		

**Table 3.** Diagnostic comparison of *Chrysosplenium guangxiense* and *C. hydrocotylifolium*.

Characters	C. guangxiense	C. hydrocotylifolium
Basal leaves and cauline leaves	Isophyllous	heterophyllous
Basal leaves	long elliptic, margin 14–24-crenate; base long cuneate	orbicular, margin 34–39-crenate; base reniform

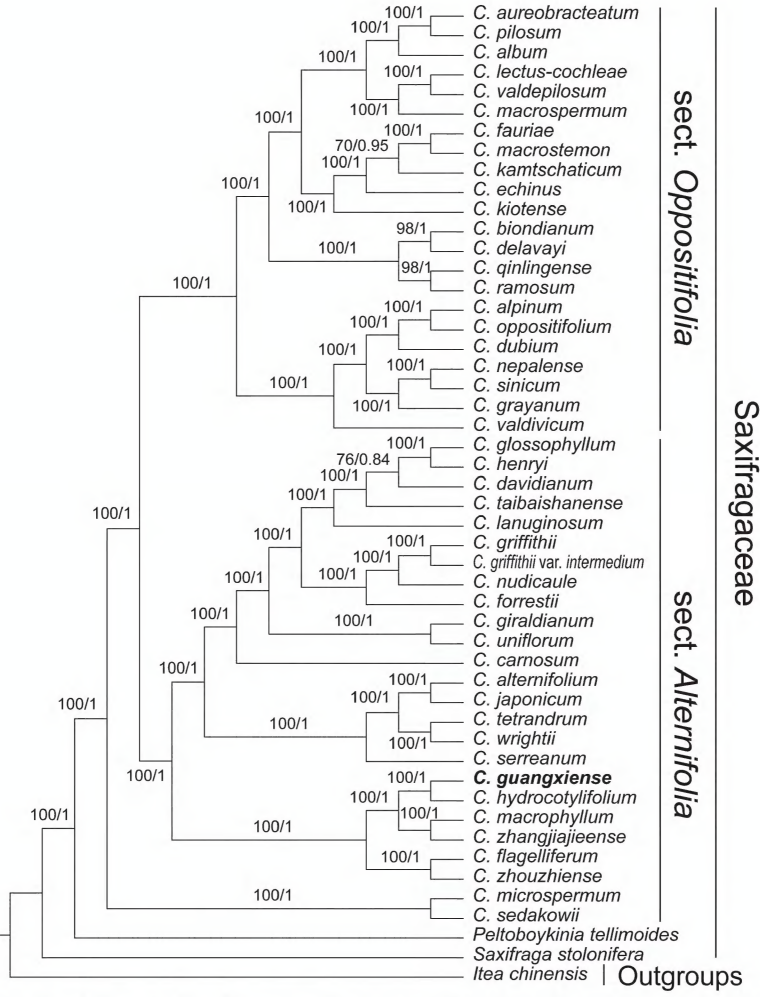


Figure 2. Phylogenetic tree of *Chrysosplenium* generated from maximum likelihood (ML) and Bayesian inference of cpPCGs+nrDNA dataset. Numbers below the branches indicate bootstrap values ( $\geq$ 50%) of the ML analyses and the posterior probability ( $\geq$ 0.5) of Bayesian analyses.

#### **Discussion**

Our phylogenetic result supported the monophyly of *Chrysosplenium* (Soltis et al. 2001; Fu et al. 2021; Yang et al. 2023). Besides two well-defined clades (denoted as sect. *Oppositifolia* and sect. *Alternifolia*), our result also revealed a third clade comprising two species from sect. *Alternifolia*, the topology of which is consistent with the previous study (Yang et al. 2023). This phylogenetic relationship indicated a non-monophyletic status of sect. *Alternifolia* and suggested that a deeper morphological character evolution across this phylogenetic framework is needed to evaluate the phylogenetic informativeness of characters.

Table 4. Summary of the complete plastid genome of Chrysosplenium guangxiense.

<b>Characteristic</b>	Chrysosplenium guangxiense	
Size (base pair, bp)	154,004	
LSC length (bp)	80,646	
SSC length (bp)	17,118	
IR length (bp)	28,120	
Number of genes	111	
Protein-coding genes	78	
rRNA genes	4	
tRNA genes	29	
GC content	37.51%	

**Table 5.** The gene statistics of the plastid genome of *Chrysosplenium guangxiense*. [Genes with one or two introns are indicated by one (\*) or two asterisks (\*\*), respectively. Genes in the IR regions are followed by the (×2) symbol].

Group of Genes	Gene Name	Number
tRNA genes	trnK-UUU, trnQ-UUG, trnS-GCU, trnG-GCC, trnR-UCU, trnC-GCA, trnD-GUC, trnY-GUA, trnE-UUC, trnT-GGU, trnS-UGA, trnS-CGA, trnfM-CAU, trnS-GGA, trnT-UGU, trnL-UAA*, trnF-GAA, trnV-UAC*, trnM-CAU, trnW-CCA, trnP-UGG, trnR-ACG(×2), trnN-GUU(×2), trnI-GAU*(×2), trnA-UGC*(×2), trnL-UAG, trnI-CAU(×2), trnL-CAA(×2), trnV-GAC(×2)	29
rRNA genes	rrn16(×2), rrn23(×2), rrn4.5(×2), rrn5(×2)	4
Ribosomal small subunit	rpsl6, rps2, rps14, rps4, rps18, rps12(×2), rps11, rps8, rps3(×2), rps19(×2), rps7(×2), rps15	12
Ribosomal Large subunit	rpl33, rpl20, rpl36, rpl14, rpl16, rpl22(×2), rpl2(×2), rpl23(×2)	8
DNA-dependent RNA polymerase	rpoC2, rpoC1*, rpoB, rpoA	4
Photosystem	psaB, psaA, psaI, psaC	5
Large subunit of rubisco	rbcL	1
Photosystem	psbA, psbK, psbI, psbM, psbD, psbC, psbZ, psbJ, psbL, psbF, psbE, psbB, psbT, psbN, psbH	15
NADH dehydrogenase	ndhJ, ndhK, ndhC, ndhB*(×2), ndhF, ndhD, ndhE, ndhG, ndhI, ndhA*, ndhH	11
Cytochrome b/f complex	petN, petA, petL, petG, petB, petD	6
ATP synthase	atpA, atpF*, atpH, atpI, atpE, atpB	6
Maturase	matK	1
Subunit of acetyl-CoA carboxylase	accD*	1
Envelope membrane protein	cemA	1
Protease	clpP**	1
Translational initiation factor	infA	1
c-type cytochrome synthesis	ccsA	1
Conserved open reading frames(ycf)	ycf3**, ycf4, ycf2(×2), ycf1(×2)	4

In our phylogenetic tree, *Chrysosplenium guangxiense* was recovered as a member of sect. *Alternifolia*, most closely related to *Chrysosplenium hydrocotylifolium* (BS = 100%, PP = 1) but had a distant relationship with *C. glossophyllum*. It was easy to distinguish *C. guangxiense* from *C. hydrocotylifolium* by the long elliptic leaves and long cuneate leaf bases (Table 3). Our morphological comparison between *C. guangxiense* and *C. glossophyllum* also showed a suite of characters, including having a robust rhizome, basal leaves with a long cuneate base and fewer teeth in margin, and larger seeds in *C. guangxiense* which helped distinguish it from *C. glossophyllum* (Table 2). Furthermore, there was a typical viviparous phenomenon of *C. guangxiense*; the mature seeds were able to germinate directly in the opening capsule (Figs 3E, 4G, H). This feature has not been reported in any other *Chrysosplenium* species so far. Therefore, our molecular and morphological evidence supports *C. guangxiense* as a distinct species that resurrected from *C. glossophyllum*. We presented the following detailed taxonomic treatment for *C. guangxiense*.

#### **Taxonomic treatment**

Chrysosplenium guangxiense H.G.Ye & Gui C.Zhang in Acta Bot. Austro Sin. 9: 57, f. 1 (1994)

Figs 3, 4

Type. Tian Lin, 11 Oct. 1989, South China Exped. 2458 (holotype: IBSC!; isotype: IBSC!). **Description.** Perennial herbs, 5.5–17 cm high. *Root* fibrous and robust. Long creeping *rhizome* developed, thick, crossed and nodular, 1-2 cm between each node, without stolons and bulbs. Flowering stem(s) always 1, erect, branched, 10-17 cm high, sparsely pilose, green, squared. Sterile branches arise from all basal leaves. Isophyllous, Basal leaves 3-6, alternate and clustered; petiole 1-6.5 cm long, white pilose; leaf blade long elliptic,  $2.2-10.3 \times 1.8-3.3$  cm, abaxially glabrous, light green, adaxially with sparse long hispid, dark green, apex rounded, margin 14-25-crenate, base long cuneate. Cauline leaves always 1, petiole 1.3-2.2 cm long; blade 2.2-4.0 × 1.2-1.9 cm, long elliptic, glabrous in the abaxial side and with sparse hispid in the adaxial side; apex obtuse; margin obtusely dentate (9-13 teeth); base broadly cuneate; veins obvious in adaxial. **Pleiochasium** 9-16 cm wide, 10-15 cm high, extremely diffused, with 5-20-flowered cyme, branches sparsely hispid, surrounded by bracts; bracteal leaves green, elliptic to broadly ovate or round, glabrous. Flowers tetramerous, actinomorphic; sepals 4 (2 pairs), flat, green, 0.9-1.2 × 2.1-4.3 mm, broadly ovate, apex acute, with margin curls outward in fruiting time; disk obvious; stamens 8, ca. 0.2 mm long, shorter than sepals; anthers orange, 2-locular, longitudinally dehiscent; ova-

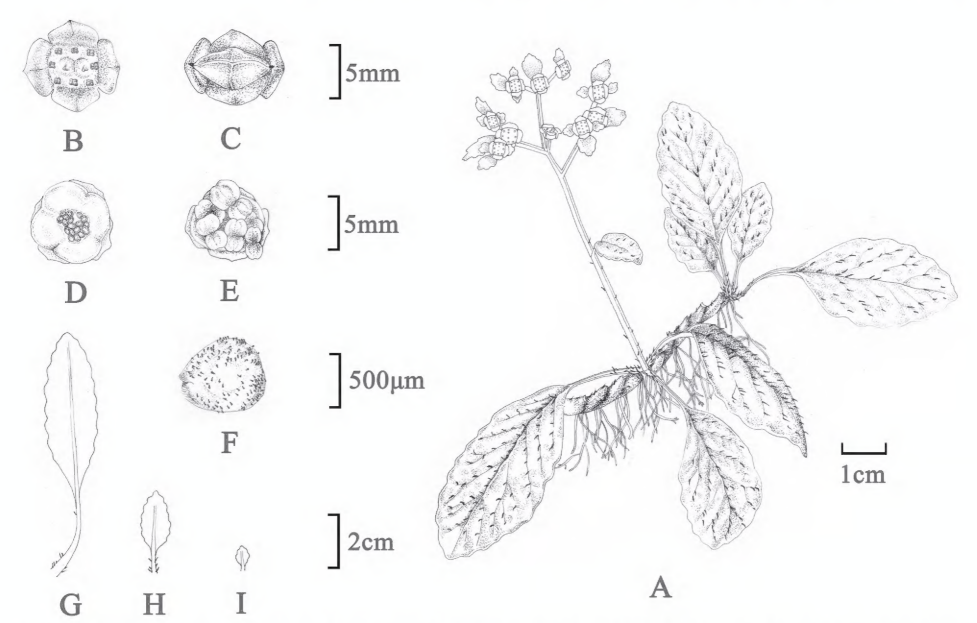
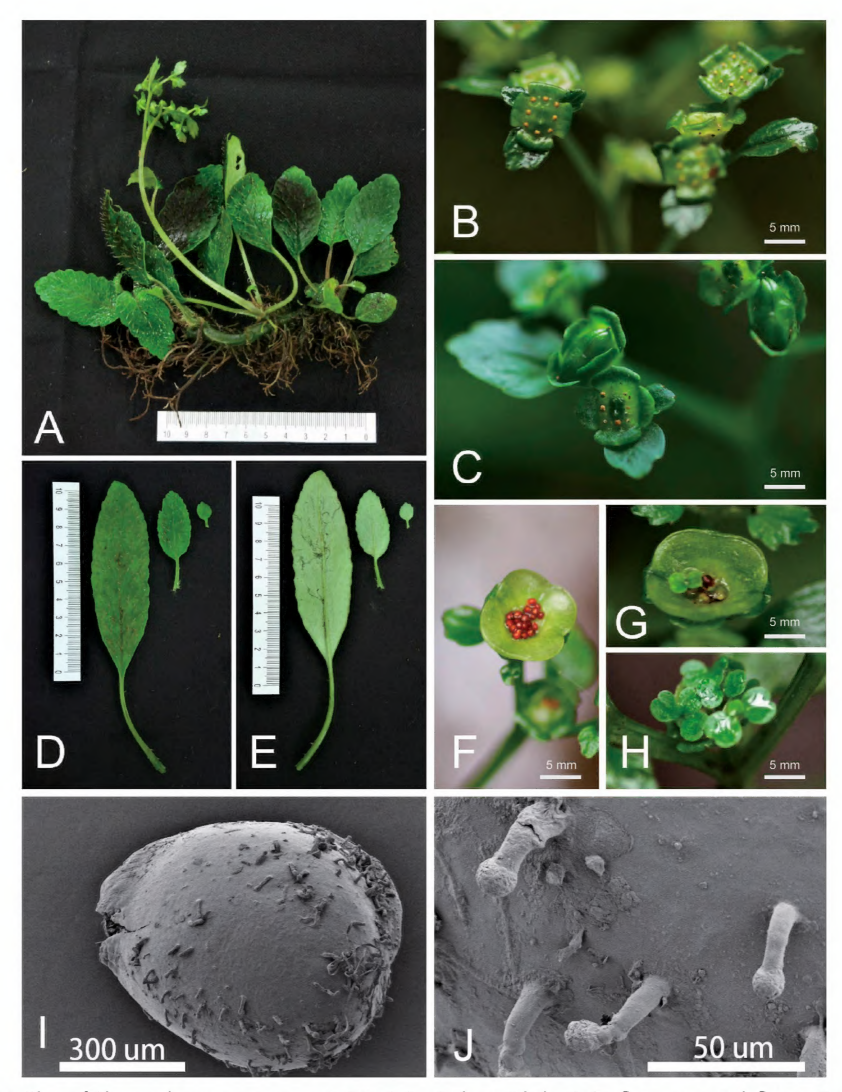


Figure 3. Illustration of *Chrysosplenium guangxiense* H.G.Ye & Gui C.Zhang A habit in flowering phase B flower C indehiscent capsule D dehiscent capsule and seeds E germinated seeds in capsule F seeds G caulline leaf H, I bracteal leaf.3



**Figure 4.** Plate of *Chrysosplenium guangxiense* H.G.Ye & Gui C.Zhang **A** habit **B**, **C** inflorescence with flowers **D**, **E** basal leaves **F** fruit and seed **G**, **H** germinated seeds and seedlings in capsule **I**, **J** SEM of seed (Photos by Hong Liu).

ry 2-locular, semi-inferior; stigma 2; styles erect, ca. 0.2 mm long. *Fruit* a capsule, 5–7 mm long, green, smooth, 2-lobed (horn-shaped), equal, dehiscent along the adaxial suture; seeds numerous, red or reddish brown, obovoid, a raphe on one side,  $594.19-855.33 \times 475.41-625.7 \mu m$ , long papillose. Viviparous.

Additional specimen examined. *Chrysosplenium glossophyllum*. CHINA. Sichuan: Kuan County (Dujiangyan City), 19 April 1930, F. T. Wang 20553 (PE!, NAS!); same locality, 6 May 1987, Xintang Ma & Zhilong Zhao 87-0521 (WCSBG!); same locality, 15 April 2013, LiXJ 353 (KUN!); same locality, 24 May 2016, Hong

Liu, HSN06644 (HSN!); same locality, 26 October 2017, Hong Liu, HSN08105 (HSN!). *Chrysosplenium guangxiense*. CHINA. Guangxi: Lingyun County, Baise City, 6 March 2014, Lingyun team 451027140305005 (GXMG!); Tianlin County, Baise City, 27 November 2019, Hong Liu HSN13356 (HSN!).

Conservation status. Chrysosplenium guangxiense is only known from two localities (IUCN criterion D2). At these two localities, the populations included ca. 200 mature individuals (IUCN criterion D1) growing in several patches. Using the IUCN methodology, C. guangxiense is classified as Vulnerable (VU) based on criteria D1 and D2: population size and the number of locations, combined with a plausible future threat that could drive this taxon to Critically Endangered or Extinct in a very short time. However, the vivipary of C. guangxiense may strengthen its adaptability to cope with future climate and environmental changes. The future threat is mainly due to grazing.

#### **Conclusions**

The phylogenetic analyses using plastomes and nuclear gene sequences of *Chrysosplenium guangxiense* reveal that *C. guangxiense* belongs to the sect. *Alternifolia*, is closely related to *Chrysosplenium hydrocotylifolium*, but distant from *C. glossophyllum* based on leaf morphology and other traits. Our findings support the resurrection of *C. guangxiense* as a distinct species and provide a detailed taxonomic treatment for its identification. The phylogenetic analyses confirm the monophyly of *Chrysosplenium* and reveal a non-monophyletic status of sect. *Alternifolia*. Further systematic studies of *Chrysosplenium* should focus on finding additional morphological characters with phylogenetic informativeness to disentangle the non-monophyletic sect. *Alternifolia*, and propose a new infrageneric classification and provide a stable framework for answering broader questions in evolutionary biology.

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#### **Additional information**

#### **Conflict of interest**

The authors have declared that no competing interests exist.

#### **Ethical statement**

No ethical statement was reported.

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#### **Author contributions**

Data curation: TGY. Funding acquisition: LH, LFF. Investigation: HL, DQL, XTC. Project administration: HL, LFF. Writing – original draft: LFF, TGY, HL. Writing – review and editing: LFF, TGY, HL.

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#### **Data availability**

All of the data that support the findings of this study are available in the main text or Supplementary Information.

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#### **Supplementary material 1**

### Phylogenetic tree of *Chrysosplenium* generated from maximum likelihood (ML) of cpPCGs dataset

Authors: Long-Fei Fu, Tian-Ge Yang, De-Qing Lan, Xi-Tang Chen, Hong Liu

Data type: jpg

Explanation note: Numbers on the branches indicate bootstrap values (left) of the ML analyses and the posterior probability (right) of Bayesian analyses.

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Link: https://doi.org/10.3897/phytokeys.243.125742.suppl1

#### **Supplementary material 2**

## Phylogenetic tree of *Chrysosplenium* generated from maximum likelihood (ML) of nrDNA dataset

Authors: Long-Fei Fu, Tian-Ge Yang, De-Qing Lan, Xi-Tang Chen, Hong Liu

Data type: jpg

Explanation note: Numbers on the branches indicate bootstrap values (left) of the ML analyses and the posterior probability (right) of Bayesian analyses.

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